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What can inactivity (in its various forms) reveal about affective states in non-human animals? A review

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Abstract: Captive / domestic animals are often described as inactive, with the implicit or explicit implication that this high level of inactivity is a welfare problem. Conversely, not being inactive enough may also indicate or cause poor welfare. In humans, too much inactivity can certainly be associated with either negative or positive affective states. In non-human animals, however, the affective states associated with elevated or suppressed levels of inactivity are still not well understood.

Part of the complexity is due to the fact that there are many different forms of inactivity, each likely associated with very different affective states. This paper has two aims. One is to identify specific forms of inactivity that can be used as indicators of specific affective states in animals. The other is to identify issues that need to be resolved before we could validly use the remaining, not yet validated forms of inactivity as indicators of affective state.

We briefly discuss how inactivity is defined and assessed in the literature, and then how inactivity in its various forms relates to affective (either negative or positive) states in animals, basing our reasoning on linguistic reports of affective states collected from humans displaying inactivity phenotypically similar to that displayed by animals in similar situations, and, when possible, on pharmacological validation. Specific forms of inactivity expressed in response to perceived threats (freezing, tonic immobility, and hiding) appear to be, to date, the best-validated indicators of specific affective states in animals. We also identify a number of specific forms of inactivity likely to reflect either negative (associated with ill-health, boredom-like, and depression-like conditions), or positive states (e.g. 'sun-basking', post-consummatory inactivity), although further research is warranted before we could use those forms as indicators of the affective states. We further discuss the relationship between increased inactivity and affective states by presenting misleading situations likely to yield wrong conclusions. We conclude that more attention should be paid to inactivity in animal welfare studies: specific forms of inactivity identified in this paper are, or have the potential to be, useful indicators of affective (welfare) states in animals.

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Highlights

Affective states linked with altered inactivity are poorly understood in animals.

Different forms of inactivity are likely to relate to different affective states.

Some specific forms (e.g. freezing) reliably indicate negative affective states.

Other specific forms are *likely to be* linked with either negative or positive states.

We propose further research directions to further validate those forms of inactivity.

What can inactivity (in its various forms) reveal about affective states in non-human animals? A review

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Abstract

Captive / domestic animals are often described as inactive, with the implicit or explicit implication that this high level of inactivity is a welfare problem. Conversely, not being inactive enough may also indicate or cause poor welfare. In humans, too much inactivity can certainly be associated with either negative or positive affective states. In non-human animals, however, the affective states associated with elevated or suppressed levels of inactivity are still not well understood.

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42

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1. Introduction

Captive or domestic animals are often described as inactive, with the implicit (*e.g.* Broom, 1988), or explicit implication that this high level of inactivity is a welfare problem (*e.g.* Zanella et al., 1996; McPhee and Carlstead, 2010). Conversely, not being inactive *enough*, for instance when external demands require increased efforts to cope with challenges and when severely sleep deprived, may also indicate or cause poor welfare; it can have dramatic adverse consequences for organisms (*e.g.* Ferrara and De Gennaro, 2001; Maslach et al., 2001). However, the affective states -- our main focus with regards to welfare¹ -- associated with these elevated or suppressed levels of inactivity are still not well understood in non-human animals (henceforth ‘animals’). This is in part because inactivity has rarely been the focus of behavioural studies and is often considered simply a default state rather than a true ‘behaviour’ (see *e.g.* Lima et al., 2005; Levitis et al., 2009).

In humans, too much inactivity can be associated with negative affective states (*e.g.* psychological distress, Muhsen et al., 2010). In animals, too, inactivity is elevated (and activity decreased) in a variety of situations where welfare is believed to be poor. In an extreme example, monkeys separated from their mothers after birth and raised alone in bare wire cages ‘sit in their home cages and stare fixedly into space’ (Harlow and Harlow, 1962). Juveniles raised in total isolation until 8 months of age needed as long as 12 to 27 weeks to begin to move at all, when placed with social companions in a room designed to trigger play (Harlow and Harlow, 1962; see Konrad and Bagshaw, 1970 for similar results in cats). Male rats exposed to social defeat become more inactive and less exploratory in novel environments than non-defeated controls (Meerlo et al., 1996a; Meerlo et al., 1996b), while buffalo cows housed in restrictive, high stocking density conditions spend more time in

¹ Defining animal welfare is a complex issue, on which not everyone agrees (see *e.g.* Fraser, 2008). Measures related to affect have however often been raised as relevant measures for assessing animal welfare; we, following *e.g.* Duncan, 2005, are principally concerned about this aspect of animal welfare.

inactive ‘idling’ than their counterparts with free access to a large outdoor yard with wallows and grazing opportunities (Tripaldi et al., 2004).

However, inactivity is also elevated in a variety of situations where welfare is self-reported (in humans, *e.g.* when receiving a gentle massage, Goats, 1994), or believed, to be good. Animals are often inactive when in familiar, safe environments where all immediate needs are met (*e.g.* Cockram, 2004; Wells, 2005; Nowak, 2006). Meanwhile, frustration of motivations to perform specific activities (deprivation: cf. Dawkins, 1988) tends to increase locomotor activity and to induce stereotypic behaviour, escape attempts and other ‘restless’ behaviour, thus likely reducing time spent inactive (*e.g.* migratory birds when caged: Mewaldt and Rose, 1960; laying hens unable to nest: Duncan, 1970; mink blocked from swimming water: Vinke et al., 2005; feed-restricted calves: Vieira et al., 2008, mink: Bildsoe et al., 1991; Hansen and Moller, 2008, horses: Benhajali et al., 2008 and rats: Prescott, 1970).

As illustrated in the examples above, the relationship between inactivity and welfare states is far from straightforward. Part of this complexity is due to the fact that inactivity is not a homogeneous category: there are many different forms, expressed in different contexts, and each likely associated with very different affective states. This paper therefore has two aims. One is to identify specific forms of inactivity that can be used as indicators of specific affective states in animals. The second is, for those specific forms of inactivity that are currently not validated as indicators, to identify issues that need to be resolved before we could validly use them as indicators of affective state. It is, however, not our goal here to provide an exhaustive list of specific examples of inactivity being associated with affective states. Instead, we intend to illustrate a rationale and to discuss further research suggestions to achieve a better understanding of the relationship between inactivity and affective states. Neither is our goal to argue that inactivity should be considered as the sole and/or ‘gold standard’ indicator of affective states in animals, but rather to highlight the potential it has,

when considered in its specific forms together with contextual information, to help infer animals' affective states. This would be valuable in a wide range of studies, such as those assessing the impact of housing, management, and other procedures on affective states, and to provide practical recommendations for eliciting positive affective states in domestic and captive animals.

In animals, linguistic self-reports on specific affective states, *i.e.* the 'gold standard' method to capture conscious affective experiences, are obviously not attainable (discussed in *e.g.* Mendl et al., 2010). Behaviours are thus the only attainable, albeit indirect, measure of putative affective states associated with a given situation. We argue that a partial solution to this problem could be to rely on our own species' experiences (providing face validity and, for some of the states, cf. 3.2.2., etiological validity, *e.g.* Belzung and Lemoine, 2011). Indeed, acknowledging that we cannot be sure what an inactive dog's, rat's or any other animal's affective experiences are, we can still make reasonable assumptions if animals' behavioural patterns in avoided (hence putatively perceived as aversive) situations are similar to those reported by humans feeling *e.g.* fearful or depressed in similarly aversive situations (*e.g.* Mendl et al., 2010). For instance, both a rat exposed to a cat (predator) (Dielenberg and McGregor, 1999), and someone frightened by a stranger entering his/her home late at night (Blanchard et al., 2001), might stay still in a location where s/he is protected behind something. We would argue that the hiding rat is in such a case likely to *feel afraid* of the perceived threat just as the motionless hiding person does. Following the same rationale, animals' behavioural patterns that are expressed in preferred or positive situations and are similar to those exhibited by humans in similar situations where they report feeling positively-valenced states, are likely to reflect positive affective states. For instance, lying in the sun, which does have a hedonic component in humans (*e.g.* Dhaenen, 1996), is likely to also be pleasurable in a diurnal rodent such as a striped mouse that chooses to lie down in the

sun when risks of predation are low (Schradin et al., 2007). Beyond analogies with humans self-reporting their feelings, additional evidence to support inferences regarding the affective valence of inactive animals' states can also come from pharmacological corroboration, *e.g.* testing that a specific form of inactivity is reduced by anxiolytics and/or increased by anxiogenics if it is believed to reflect fear. This provides evidence of construct validity and, in cases where a range of treatments was tested, discriminant validity (see *e.g.* Cronbach and Meehl, 1955). In this review, we briefly discuss how inactivity is defined and assessed in the literature, and then discuss how it relates to affective (welfare) states. In humans, affective states can be categorized in terms of two fundamental underlying dimensions: the valence – *i.e.* whether the experience is perceived as negative or positive, punishing or rewarding, unpleasant or pleasant– and the reported activation – *i.e.* high or low arousal (*e.g.* Russell and Barrett, 1999). Theoretical and empirical studies (reviewed in Mendl et al., 2010) suggest that negative high-arousal affective states (*e.g.* feeling fearful) are principally associated with perceiving and reacting to threats or dangers, while negative low-arousal states (*e.g.* feeling sad) are likely to be associated with experiences of loss or lack of reward, and may promote low activity and energy conservation when resources are lacking. Positive high-arousal affective states (*e.g.* excitement) are likely to be associated with appetitive motivational states, and function to facilitate seeking and obtaining rewards; while positive low-arousal affective states (*e.g.* calmness) are instead expected to be associated with low levels of experienced threat, perhaps facilitating the expression of maintenance activities. In this paper, we rely on this framework in order to categorize affective states, and those specific forms of inactivity associated with such states. This classification of specific forms of inactivity displayed by animals will be supported by linguistic self-reports collected in humans displaying phenotypically similar inactivity in similar situations, and, when possible, by pharmacological validation.

2. How is inactivity defined and measured?

Although inactivity may seem fairly straightforward to define, there is some variation in exactly what the term encompasses. In some cases, the hypotheses under test relate to specific forms of inactivity (*e.g.* in rats: sleeping, or lying non-alert with both eyes closed: Abou-Ismaïl et al., 2008; freezing, a complete absence of visible movement except breathing: Fanselow, 1982), and so only those forms are assessed. However, in other cases, its operational definition seems to be a by-product of the methods used to assess activity. This usually occurs when inactivity is not the main focus of the study but is measured simply as a part of the time budget or to control for ‘activity’ levels because they influence the variable of interest. When it is studied in free-living animals, activity is usually assessed using radio-tracking or similar technology, and so any time not in locomotion is called inactivity. Some studies of captive animals similarly equate ‘activity’ with locomotion. Laboratory animal research, for example, often infers activity levels from proxy measures such as the number of entries into closed arms (and therefore locomotion) in the elevated plus maze, a test of anxiety (*e.g.* Louvart et al., 2005), while some agricultural studies use pedometers (*e.g.* O’Callaghan et al., 2003). Therefore, inactivity in those studies would include time spent stationary but performing purposeful movements such as grooming. However, in applied ethology research using video or live observation, any movement is typically considered activity, even if the animal remains in one place; for example, kicking (*e.g.* Rushen et al., 2001) and eating (*e.g.* Rochlitz et al., 1998; Burrell and Altman, 2006) are not categorized as inactive. Thus, the most common definition of inactivity is being relatively motionless, and although it is rarely stated explicitly, this means no movement with an apparent function (*e.g.* grazing or chewing a bite of food) but would include other slight movements (*e.g.* turning the head or shifting positions). Finally, research on anticipatory behaviour sometimes discusses

hyperactivity or decreases in activity during anticipation, but actually measures the number of behavioural transitions rather than total amount of time spent moving or stationary (*e.g.* van den Bos et al., 2003). Studies using any of these definitions of inactivity will be discussed here.

3. Increased inactivity / decreased activity as a sign of poor affective states

Increased inactivity likely to be associated with negative affective states will be considered here, distinguishing specific forms of inactivity expressed in response to a perceived threat (likely to be associated with negative high-arousal affective states, Mendl et al., 2010, cf. Introduction) from those expressed in situations where threat levels but also chances of getting rewards are low (likely to be associated with negative low-arousal affective states).

3.1. Increased inactivity / decreased activity displayed in response to a perceived threat

Inactivity that may occur in response to a perceived (frightening) threat stimulus will be presented here, targeting more specifically the freezing, tonic immobility and hiding responses.

Freezing (sometimes also termed *attentive immobility*) is a common response in the face of an immediate (perceived) threat in various species, where an individual becomes abruptly motionless, monitoring a perceived source of danger (reviewed by Boissy, 1995). A ‘freezing-like’ state, characterised by increased immobility and rigidity (quantified using a force platform) and a reduced heart rate (also termed ‘fear bradycardia’), has been described in humans. This state has been observed in healthy people viewing pictures of mutilation (that they self-rated as negative in valence and high in arousal) compared to neutral or

positively rated pictures (Azevedo et al., 2005; Facchinetti et al., 2006), as well as in patients diagnosed with panic disorders when seeing either pictures of mutilation or images that were anxiogenic due to their pathologies (*e.g.* crowded areas for agoraphobic patients), compared to neutral pictures (Lopes et al., 2009).

In animals, freezing is used to quantify fearfulness in many behavioural tests (see *e.g.* Bouton and Bolles, 1980; Forkman et al., 2007). It has been particularly well-described in rodents, on which we will focus here, and can be operationally defined as ‘the absence of all visible movement of the body and vibrissae, except for movements necessitated by respiration’ (Fanselow, 1982). The description can also include (species-specific) postural elements, such as a ‘characteristic immobile, crouching posture’ in rats, and autonomic changes, *i.e.* a decreased heart rate (similarly to the ‘fear bradychardia’ recorded in freezing humans) and an increased respiratory rate (Fanselow, 1984). Freezing can be induced by exposing rats to a predator odour (*e.g.* Wallace and Rosen, 2000; Knox et al., 2012), as well as to conditioned stimuli or contexts paired with aversive experiences (electric shocks: *e.g.* Fanselow, 1984; Richmond et al., 1998; Luyten et al., 2011; carbon dioxide inducing dyspnoeic suffocation: Mongeluzi et al., 2003). The more intense the aversive stimuli are, the longer the rats remain frozen (Fanselow and Helmstetter, 1988; Wallace and Rosen, 2000; Mongeluzi et al., 2003; Santos et al., 2005, but see Leaton and Borszcz, 1985 for non-monotonic effects). Freezing is ‘not a simple suppression of activity’ (Fanselow, 1984) but a highly aroused state (Bracha, 2004): rats’ percentage of time spent freezing when exposed to a conditioned stimulus previously paired with electric shock was positively correlated with the amplitude of their acoustic startle response in the presence of that conditioned stimulus (Leaton and Borszcz, 1985). Interestingly, neither food deprivation (an aversive experience but not assumed to induce ‘fear’, Maren and Fanselow, 1998; Heiderstadt et al., 2000), nor exposure to smells unrelated to predation (*e.g.* butyric acid, banana and pear odours, Wallace

and Rosen, 2000; Knox et al., 2012) induce freezing responses. Moreover, in rats where the risk of infanticide by unrelated adult males ends around weaning, exposure to a sexually experienced, unrelated male rat induces freezing in young rats only before they reach a natural weaning age (around 26 d), whereas exposure to a cat odour (a predation risk relevant during all life stages) induces the freezing response in rats before and after weaning age (Wiedenmayer and Barr, 2001). Additional evidence that the freezing response is associated with negative valence in rodents comes from pharmacological corroboration: in rats, while freezing in response to a cat exposure was gradually suppressed over repeated daily exposure (Farook et al., 2004), administration of anxiogenic drugs restores the freezing behaviour. Conversely, the administration of anxiolytic drugs (*e.g.* midazolam, diazepam) reduces the duration of rats' place-conditioned freezing response (Fanselow and Helmstetter, 1988; Verleye and Gillardin, 2004; Santos et al., 2005).

Another specific form of inactivity also displayed in response to a threat in various species is **tonic immobility (TI)** (*e.g.* Gallup et al., 1971b). In humans, 'TI-like' states are characterised by a temporary behavioural state of motor inhibition, associated with tremors, eye closure, increased breathing, and coldness, and have been reported to occur in response to situations involving intense fear and physical restraint such as interpersonal trauma (sexual assault, torture, armed robbery)² (Galliano et al., 1993; Abrams et al., 2009). Although such studies are non-experimental in nature (and therefore results may not be generalised to any human population), TI-like responses appear to be far from uncommon. For example, across several studies reviewed in Galliano et al. (1993), from 12% to 50% of the studied victims of rape/sexual assaults were 'paralyzed' motionless and did not resist their attackers in any way. Displaying TI-like responses during traumatic episodes has also been reported to positively

² Although less studied, intense fear associated with accident-related trauma and the unexpected death of a loved one have also been reported to sometimes induce a TI-like state in humans despite not involving physical restraint.

241 correlate with longer-term psychological impairments such as depression, anxiety and Post-
242 Traumatic Stress Disorder (Abrams et al., 2009; Volchan et al., 2011). In laboratory studies,
243 ‘standing still (paralyzed with fear)’ is one of the defensive strategies subjects predicted they
244 would display in response to threat scenarios involving nearby threat stimuli and
245 inescapability of the threat/situation, such as ‘*Late at night you are alone in an elevator.*
246 *When it stops and the doors open, a rough looking stranger gets in fast to attack you,*
247 *blocking your exit*’ (Blanchard et al., 2001; Shuhama et al., 2008).

248 In animals, TI has been reported in several taxa (see *e.g.* Forkman et al., 2007), and is
249 particularly well-described in birds, where it has been defined as a ‘reversible state of
250 (‘catatonic-like’) profound motor inactivity following brief exposure to physical restraint
251 (*e.g.* 15s), which may last from a few seconds to over several hours’ (*e.g.* Gallup et al.,
252 1971b). It is also characterised by a suppression of vocalisation, as well as TI-specific (not
253 observed in freezing) muscle tremors in the extremities and intermittent eye closures. As for
254 freezing, physiological correlates include bradycardia and increased respiratory rate, as well
255 as a TI-specific (not reported in freezing) decrease in body temperature (*e.g.* Gallup et al.,
256 1971b; Nash et al., 1976). TI is an aroused state: electroencephalographic activity in animals
257 displaying TI has been reported to be often the same as that of waking animals (*e.g.* in
258 rabbits: Klemm, 1966; in opossum *Didelphis virginiana*: Barratt, 1965, review in Gallup,
259 1974; Whishaw et al., 1982). A ‘TI-like’ response has also been described in certain domestic
260 goats, known as ‘fainting’ goats becoming ‘perfectly rigid when suddenly surprised or
261 frightened’ (Lush, 1930). This response is caused by a hereditary genetic disorder (*congenital*
262 *myotonia*, Clark, 1939), although physiological correlates of this state have not been
263 investigated. In birds (chicks unless otherwise specified), a variety of aversive manipulations
264 before TI induction increase the TI duration and/or propensity of the bird to display the TI
265 state, including exposure to electric shocks or to conditioned stimulus signalling shocks

(Gallup et al., 1970; Gallup, 1973), rough handling (bird inverted for 30s: laying hens and broilers, Jones, 1992), and exposure to loud noise (Gallup et al., 1970). While freezing in rodents appears to be a risk-assessment behaviour to a (perceived) distant threat (Blanchard et al., 2011), TI happens *following physical restraint*, and has been suggested to be an anti-predation response even after the animal has been captured. Such ‘death-feigning’ might induce the predator to loosen its hold (Gilman et al., 1950; Engel and Schmale, 1972; Sargeant and Eberhardt, 1975; see Thompson et al., 1981 for evidence TI can deter predators). This response seems specific to fear-inducing situations: food-depriving chickens, which is aversive but not likely frightening, does not increase these animals’ TI duration (Gallup and Williamson, 1972). Additional evidence supporting the negative valence and high arousal of the TI response in birds comes from pharmacological validation: in chicks, pre-TI-induction administration of adrenaline (Braud and Ginsburg, 1973) and corticosterone (Jones et al., 1988) increases the TI duration and/or the propensity of birds to display TI, while a pre-TI-induction tranquilizer injection reduces the duration of the TI response (Gallup et al., 1971a).

Another -- perhaps less species-specific -- form of inactivity that can be displayed in response to a perceived threat is **hiding**. In humans, hiding (protecting oneself behind something) is one of the defensive strategies chosen by subjects in laboratory studies in response to fearful threat scenarios such as ‘*Late at night... you are sleeping alone in your bed. You suddenly wake up feeling that you heard a suspicious noise*’. Not surprisingly, the presence of a place of concealment or protection in the scenario promoted the hiding choice; so did distant (rather than close) threat stimuli (Blanchard et al., 2001; Shuhama et al., 2008).

In animals, hiding can be defined operationally using location (provided that there are locations suitable for hiding in the environment): hiding animals are ‘remaining stationary

and out of sight or camouflaged using any kind of shelter or visual barrier' (Meagher et al., 2013). In rodents, exposure to a predator or to its odour initially induces a hiding response (e.g. rats exposed to a worn cat collar: Dielenberg and McGregor, 1999; mice repeatedly exposed to a rat moving around on top of their cages: Dalm et al., 2009). In laboratory cats, exposure to complex stressors (involving unpredictable mildly aversive procedures) increases time spent awake/alert and attempting to hide, and suppresses active exploratory and play behaviour (Carlstead et al., 1993b). Translocation to novel environments also induces hiding in felids (leopard cats: Carlstead et al., 1993a; quarantined domestic cats: Rochlitz et al., 1998). There is also pharmacological evidence to help infer the negative valence and high arousal of the hiding response: in rats, anxiolytic administration (the benzodiazepine drug midazolam) reverses rats' hiding response to a worn cat collar (Dielenberg and McGregor, 1999), and increases the proportion of time spent exploring in open arms in an elevated plus maze, while anxiogenic substances (e.g. caffeine) increase the time spent hiding in the closed arms of the maze (Pellow et al., 1985).

Freezing, TI and hiding are specific forms of inactivity expressed in response to a (perceived) actual or potential threat, both in humans and in animals. In both, they are reduced by anxiolytics and increased by anxiogenic drugs. Freezing, TI and hiding therefore appear to be valid indicators of a negative, highly aroused affective state, and to date, our best examples that specific forms of inactivity can be used as trustworthy indicators of specific affective states (in this case, 'fear-like' states) in animals.

3.2. Increased inactivity / decreased activity likely to be associated with negative low-arousal affective states

Increased inactivity expressed in situations where both threat levels and chances of getting rewards are low (likely to be associated with low arousal negative affective states: Mendl et al., 2010, see Introduction) will be discussed here, specifically targeting sickness, depression-like, and boredom-like conditions.

3.2.1. Inactivity and ill-health

Lethargy (*i.e.* a state of decreased mental activity, characterised by sluggishness, drowsiness, inactivity, and reduced alertness, APA, 2013) is a well-established component of sickness behaviour, which is the ‘coordinate set of subjective, behavioural and physiological changes that develop in sick individuals during the course of an infection’ (Dantzer, 2004). ‘Sick individuals are somewhat depressed and lethargic’ and ‘show little interest in their surroundings and stop eating and drinking’ (Dantzer, 2004). Reduced activity here is considered a strategy of energy conservation in order to allow the full development of a fever (which is associated with and plays a critical role in recovery from many pathogenic infections), and so has an eventual benefit. However, this inactivity is very likely linked with negative affective states, as it is ‘very often accompanied by pain’ (Dantzer, 2004), and in humans, a transient depressive state has been reported to occur as an infectious episode develops (Aubert, 1999). Sickness behaviour is common to many mammalian species (Hart, 1988; Maes et al., 2012): ‘lethargy’, ‘listlessness and disinterest in social interactions with the environment’, ‘behavioural inhibition’, and ‘reduction of locomotor activity, exploration and grooming’ have also been observed in sick animals. For instance, rats challenged with bacterial and viral mimetics show decreased voluntary running wheel activity and, broadly, less movement in their home cage (Hopwood et al., 2009). The general decrease in behavioural activities in sick animals has been shown to reflect changes in motivational state rather than a simple consequence of weakness: for example, if pups are removed from the

nest of lactating mice whose behavioural activity is depressed by LPS injection, the sick mothers interrupt their sickness behaviour to bring the pups back to the nest, then return to inactive recuperative behaviour (Aubert, 1999). As in humans, lethargy in sick animals is likely to be associated with negative affect, such as pain (*e.g.* in dogs: Wiseman et al., 2001).

Perhaps more broadly, ill-health in humans (including not only infectious sickness but *e.g.* injury, post-operative conditions, and chronic back disorders) reduces both voluntary (*e.g.* work, recreational) and obligatory (*e.g.* self-care) activities (*e.g.* Tait et al., 1990). It seems to have the same effect in animals: poor health conditions can increase the proportion of time spent awake but lying down (*e.g.* postoperative pain in rabbits: Leach et al., 2009 and horses: Pritchett et al., 2003; ear notching and tagging in piglets: Leslie et al., 2010; lameness in dairy cattle: Chapinal et al., 2010; Calderon and Cook, 2011 and in broilers: Weeks et al., 2000), whereas analgesia reduces time lying down in lame animals (*e.g.* dairy cattle: Schulz et al., 2011; Offinger et al., 2013). Adult zebrafish (*Danio rerio*) injected with acetic acid (a noxious chemical stimulus) display decreased swimming activity (Correia et al., 2011; but see Steenbergen and Bardine, 2014 for an opposite effect on zebrafish larvae water-exposed to acetic acid). High activity of a shoal could thus indicate that its members are healthy, and joining it could be beneficial for fitness (*e.g.* active fish can be quicker to find food patches and more confusing for predators), which might be part of the reason that, although shoaling zebrafish usually prefer to join larger shoals, this preference can be shifted to a smaller shoal if its members are comparatively more active than the fishes in the larger shoal (Pritchard et al., 2001).

Poor health conditions associated with negative affective states such as pain, appear to increase inactivity, both in humans and in animals, in each of which they are reduced by analgesic drugs. Such inactivity is therefore likely to be associated with negative affective states. Poor health-induced inactivity is, however, less specifically described (*i.e.* overall

increased inactivity / decreased activity) than, for example, those forms of inactivity displayed in response to a perceived threat. The presence of signs of ill-health (*e.g.* fever, injury) and/or knowledge of specific contexts in which inactivity increases (*e.g.* post-surgery) therefore appears crucial to infer the affective state associated with such inactivity.

3.2.2. Inactivity and depression-like states

In humans, clinical depression -- by which we mean ‘major depressive disorder’ or experiencing ‘depressive episodes’, to encompass DSM-V (Diagnostic Manual of Mental Disorders fifth edition, American Psychiatric Association [APA], 2013) and ICD-10 (International Statistical Classification of Diseases and Related Health Problems, World Health Organisation [WHO], 1994) terminologies -- is a common mental illness diagnosed by the co-occurrence of several affective, cognitive and behavioural symptoms. These include a ‘depressed (low, sad) mood most of the day, nearly every day, as indicated by either subjective report (*e.g.* feels sad, empty, hopeless) or observation made by others (*e.g.* appears tearful)’ (APA, 2013, P160). A common trigger is chronic stress, such as that arising from aversive life events or chronic pain or illness (Blackburn-Munro and Blackburn-Munro, 2001; Siegrist, 2008; Hammen et al., 2009; APA, 2013). Cognitive changes can be associated with depression and may act as mediators in some subjects, being hypothesised to contribute to the onset and/or maintenance of the disease (Beck, 1967; Gotlib and Krasnoperova, 1998). One such change, ‘learned helplessness’, is proposed to occur ‘when highly desired outcomes are believed improbable or highly aversive outcomes are believed probable, and the individual comes to expect that no response in his repertoire will change their likelihood’ (Abramson et al., 1978).

With respect to inactivity, a low, sad mood may induce increased inactivity even in healthy people. For instance, Rucker and Petty (2004) showed that inducing sadness in

consumers in a laboratory setting yields a preference for an advertised product promoting passivity (a vacation resort framed as a place to relax and rest), while inducing anger yields a preference for a product promoting activity (a vacation resort framed as a place to enjoy sports and activity). Accordingly, clinically depressed patients have been reported to be more inactive -- by which we mean here a decrease in a variety of daily activities -- than their non-depressed counterparts. This includes 'not doing fun activities or chores that need to be accomplished' (Knowles, 1981), and reported difficulties initiating or completing social and non-social activities (Baker et al., 1971; Schelde, 1998; APA, 2013). Reduced physical activity (both mild, such as walking and gardening, and more vigorous, such as playing sports) has been associated with clinical depression (Seime and Vickers, 2006; Lindwall et al., 2011), while – cautiously³ - increased exercise has been reported in several reviews or meta-analyses to improve depressed mood and/or anxiety (e.g. Byrne and Byrne, 1993; Dunn et al., 2001; Seime and Vickers, 2006; Davis and Dimidjian, 2012).

Could inactive animals, or at least those displaying (certain forms of) inactivity (in certain contexts), be experiencing 'depression-like' states?⁴ Presumably yes: dogs and cats (e.g. Fox, 1968, p. 357) and elephants (Mason and Veasey, 2010) have anecdotally been suggested to become highly inactive when deprived of their owners or after the loss of a social companion, as have apes housed long-term in barren environments in laboratories or zoos (e.g. Engel, 2002, p174; Brune et al., 2006), and socially deprived monkeys (e.g. Harlow and Harlow, 1962; Harlow and Suomi, 1974; Suomi et al., 1975). Because the aetiology corresponds to

³ Due to methodological biases present in one or more of the studies included in the review / meta-analysis, e.g. people are not systematically randomly assigned to treatment groups and/or there are potential confounds or no control groups and/or the amount of physical activity applied as a treatment is based on patients' self-reporting (no verification) and/or conclusions are expanded from normal subjects to clinical samples. These biases, however, are spread across individual studies, and a variety of biological and psychological mechanisms could explain the reported benefit of exercise on mood, cautiously suggesting that this commonly report effect might be a 'trustable' phenomenon.

⁴ Discussing in detail whether or not non-humans can become *clinically depressed-like* - i.e. show states that share the same or most of the properties of those described in clinically depressed patients -- would go beyond the scope of this paper. However, even if the quality and quantity of current evidence are not yet sufficient to conclude this with certainty, both circumstantial and experimental evidence have led several authors (including us) to hypothesise that depression-like states occur in other animals as well (see e.g. Ferdowsian et al. 2011; Hennessy et al., 2014; Fureix et al., 2015).

410 theories of human depression emphasizing aversive life events and chronic stress as a
411 common trigger, such inactivity is likely to be associated with negative affect.

412 Moreover, the cognitive feature of learned helplessness has also been shown in animals,
413 and is a phenomenon typically accompanied by an overall decrease in activity (see *e.g.*
414 Mineka and Hendersen, 1985). Indeed, although the term learned helplessness referred
415 initially to a deficit in avoidance learning induced by repeated exposure to uncontrollable
416 shock (reviewed by *e.g.* Maier and Seligman, 1976), the meaning of the label has now been
417 expanded; it is sometimes applied to any ‘passive’ behaviour (*i.e.* quiescence or the absence
418 of active responses to stress, such as escape attempts; cf. Oxford English Dictionary, 2005)
419 that appears to result from exposure to uncontrollable stressors (Maier, 1984; see also
420 Wemelsfelder, 1990; Carlstead, 1996). For instance, sheep moved from pasture to
421 inescapable indoor crates (Fordham et al., 1991) and laboratory rodents placed in an
422 inescapable container filled with water (known as the Porsolt Test, reviewed in *e.g.* Deussing,
423 2006), both begin by reacting to the situation with agitation, but end up displaying inactivity
424 and unresponsiveness. According to the above-mentioned expanded definition, this eventual
425 response would reflect learned helplessness. Again, because the aetiology corresponds to
426 cognitive theories of human depression (Beck, 1967; Abramson et al., 1978; Gotlib and
427 Krasnoperova, 1998), this inactivity is believed to be a depression-like behaviour, and
428 therefore associated with a negative affective state. In mice and rats that ‘cease struggling and
429 remain floating motionless in the water, making only movements necessary to keep their head
430 above water’ (Porsolt et al., 1977) in the Porsolt test, additional support comes from the fact
431 that this specific form of inactivity is both amplified by stressors and alleviated by
432 antidepressants (Porsolt et al., 1977; Cryan et al., 2002; Matthews et al., 2005; Deussing,
433 2006; McArthur and Borsini, 2006). It also co-varies with other depression-like symptoms,
434 such as anhedonia (Strekalova et al., 2004), *i.e.* the loss of pleasure, a key feature of human

clinical depression (APA, 2013). The hypothesis that the term learned helplessness might also be applicable to captive animals that seem very passive or inactive in their home environment is also supported by findings that animals reared in socially isolated and/or barren cages are more vulnerable to developing learned helplessness in avoidance learning paradigms than those reared in more socially and physically complex, and presumably controllable, environments are (Seligman, 1972; Chourbaji et al., 2005).

Finally, Fureix and colleagues (2012, 2015) recently described long-lasting inactive ‘withdrawn’ states in certain riding horses, characterised by bouts of unresponsiveness, remaining motionless with unblinking eyes with an apparently fixed gaze (reminiscent of the reduced responsiveness and reduced interactivity of some depressed human patients) and anhedonia. These states also correlate with stereotypic behaviour (a possible marker of current, but also past exposure to stressors). While the aetiology of this specific form of inactivity is currently unknown, its association with key features of human clinical depression makes it likely to be associated with negative affect.

Do these findings demonstrate with certainty that these inactive animals are clinically depressed, in the same way as depressed patients showing decreased variety in their daily activities? The quality and quantity of current evidence are not yet sufficient to conclude this. Moreover, while some forms of inactivity are highly specific (‘floating’ in rodents, ‘withdrawn’ states in horses), others are not (*e.g.* passivity when exposed to inescapable, uncontrollable stressors). However, that inactivity appears in contexts similar to those that trigger the appearance of clinical depression in humans, or co-varies with key symptoms of this pathology (*e.g.* anhedonia) is sufficiently consistent with the hypothesis to make additional research into these topics, including how this inactivity would be modulated by anti-depressant drug treatments, very worthwhile.

3.2.3. Inactivity and boredom

Boredom is a negative affective state induced by monotony or lower-than-optimal levels of stimulation. In addition to self-report, this definitional link to negative affect is supported by evidence of high motivation to avoid the state; for example, boredom-prone people show a preference for activities that are perceived as risky and therefore frightening to most people, but that increase stimulation levels, such as bungee-jumping (Michel et al., 1997). Self-reports, however, are key to identifying boredom and situations that induce it (Harris, 2000).

In humans, lethargy is a common symptom (see *e.g.* Inglis, 1983), although this is often seen following a period of sensation-seeking (Taylor and Cohen, 1972; Inglis, 1983; cf. Berlyne, 1960 for an alternative possible time course) and thus in some cases, restlessness may be seen rather than inactivity (reviewed by Kirkden, 2000). Imposed inactivity can also be a cause of boredom (Berlyne, 1960; Heaman and Gupton, 1998). Most theoretical discussion categorizes boredom as a state of under-arousal (*e.g.* Fiske and Maddi, 1961; Stevenson, 1983; Mikulas and Vodanovich, 1993) given its association with low stimulation, although Berlyne (1960) postulated that prolonged monotony can lead to increases in arousal; others have similarly considered boredom simply as a state of ‘non-optimal’ arousal (Eastwood et al., 2012). While there is some evidence from humans that arousal may sometimes be elevated during boredom (*e.g.* EEG data from subjects exposed to sensory deprivation after sleeping as much as possible: Berlyne, 1960), other studies have found decreasing arousal over time when engaged in a boring task (*e.g.* Pattyn et al., 2008). Also supporting the association with under-arousal, and thus supporting its inclusion in this section of our discussion, methods of avoiding boredom are likely to increase arousal: these include consumption of recreational drugs (Samuels and Samuels, 1974), which commonly include stimulants (Boys et al., 2001), and participation in thrill-seeking activities, as previously mentioned.

Due to the dependence on self-reported affect for identifying boredom in humans, this state has been subject to little empirical investigation in animals, where self-report is impossible and thus affect cannot be assessed directly. However, captive animals commonly face monotonous environments, often less complex or lower in stimulation than those in which their ancestors evolved (in some cases, even those they experienced themselves early in life). For this reason, theory suggests that they would also find such situations aversive. For example, McFarland (1989) proposed that when captive animals' immediate physical needs are met but they cannot pursue other activities that would occupy their time in the wild such as reproduction or mating, they are left in a state of 'limbo' and are likely to suffer because most species will not have evolved methods of coping with such a situation. Veissier et al. (2009) also argue that since sheep are sensitive to the same features of stimuli that induce boredom in humans, they are potentially capable of experiencing it; the same argument could be applied to many species. Inactivity is generally accepted as a common consequence of housing in relatively barren cages or enclosures and interpreted as a sign of poor welfare (DeMonte and LePape, 1997), which many people attribute to boredom (*e.g.* Stevenson, 1983; Woodgush and Beilharz, 1983). Conversely, increasing activity or behavioural diversity through provision of opportunities to interact with stimuli is assumed to improve welfare (*e.g.* pigs: Woodgush and Beilharz, 1983; chimpanzees: Celli et al., 2003; dogs: Wells, 2004). Sometimes this assumption has been supported by improvements in other welfare indicators (*e.g.* Paquette and Prescott, 1988) or by animals' preference for the enrichment (*e.g.* Rozek et al., 2010).

Both the use of the term 'boredom' in animals and its relationship to inactivity still need validation, however. To provide a starting point for this work, Meagher and Mason (2012) proposed an operational definition based on motivation to obtain stimulation, which should be a universal symptom. The validity of this operational definition was supported by the fact

that this motivation was elevated in captive mink housed in non-enriched cages, predicted to experience more boredom-like states. Thus, compared to mink housed in a preferred (Dallaire et al., 2012) and more stimulus-rich environment, the mink behaved as bored humans would. This method of assessment relied on measuring activity when given an opportunity to avoid boredom rather than directly assessing inactivity in the hypothesized boredom-inducing situation, because the latter might vary with time and between individuals (as in humans), among other reasons. However, the study also identified a tentative link between the apparent boredom and a specific subtype of inactivity when undisturbed in the home cage (lying down with the eyes open). Future work could use self-administration of stimulants to further validate the concept of boredom in barren-housed animals and its association with inactivity, predicting that very inactive individuals in non-enriched cages would be most likely to self-stimulate. At least until such work has been carried out for a given species, inactivity should be used as an indicator of boredom with extreme caution: although high levels of inactivity in monotonous environments may well be associated with boredom, the alternative response of restlessness would make this indicator prone to false negatives (see also 5.1.), in which an environment that is in fact boring does not increase group-level or even individual-level inactivity levels.

4. Increased inactivity / decreased activity as a sign of good affective states

Increased inactivity or decreased activity likely to be associated with positive affective states will now be discussed, again distinguishing inactivity likely to be associated with high and low-arousal positive states.

4.1. *Increased inactivity / decreased activity likely to be associated with positive highly-aroused affective states*

As stated in the Introduction, positive, highly-aroused affective states are likely to be associated with appetitive motivational states, and function to facilitate seeking and obtaining rewards (Mendl et al., 2010). Being *inactive* in order to *favour* reward *acquisition* sounds intuitively unlikely to happen, and examples are rare even in humans, with perhaps the exception of yogi meditation, which has been self-reported by meditators to be a highly-aroused pleasant state (Cahn and Polich, 2006). Chess players close to winning a game and focused on choosing the best strategy could also perhaps experience a highly-aroused and pleasant motionless state; this is likely one example of what Csikszentmihalyi (1975; 1990) termed a “flow” state. Flow states involve being concentrated on a task that is achievable but sufficiently challenging to require focused attention and skill, and are self-reported as being enjoyable, at least in retrospect (reviewed in Csikszentmihalyi 1990). However, although the absence of evidence is not a proof of absence, one may reasonably doubt that animals practice meditation or play chess. An animal example in this category might be cats ‘stalking’: adults stalking prey (Wise, 1974) and kittens playing (Bateson and Young, 1981) temporarily restrain any movement and stay perfectly motionless. Cats have been considered to become ‘hypoactive’ while anticipating food rewards, displaying reduced behavioural transitions between the offset of a conditioned stimulus and the onset of an unconditioned stimulus in a Pavlovian conditioning paradigm (van den Bos et al., 2003). According to the authors, this might be expected as ‘they [*cats*] normally employ a ‘sit-and-wait’ tactic while close to their prey’. Bouts of immobility while stalking could therefore tentatively be seen as a (cat-specific) form of inactivity that would favour reward acquisition, but the affective state(s) associated with such a behaviour are clearly not validated yet (see *e.g.* Bassett and

Buchanan-Smith, 2007 for evidence that opposite affective states are sometimes associated with anticipation).

4.2. *Increased inactivity / decreased activity likely to be associated with positive low-arousal affective states*

Increased inactivity expressed in situations with low levels of experienced threat, and that facilitates the expression of maintenance, consolidation and recovery (cf. Introduction) will be discussed here, targeting more specifically ‘sun-basking’ and post-consummatory inactivity. Note that resting will be discussed later (see part 5.4.).

4.2.1. ‘Sun-basking’ inactivity

In humans, UV exposure activates known reward centres in the brain (Harrington and colleagues 2012, cited in Fell et al., 2014), and lying in the sun or, in other words, ‘sun-basking’, has hedonic properties (Dhaenen, 1996; Loas et al., 2000); it might even turn into an addictive behaviour (Fell et al., 2014). According to Balcombe (2009), ‘animals’ lives afford them the opportunity to experience a wealth of other pleasures beyond the realms of food, sex and touch, such as basking in the sun or seeking shade’. Supplying captive wombats (*Lasiorninus latifrons*) with feed and olfactory items (so-called enrichments, but note that such items did not reduce the time spent displaying stereotypic behaviours in this study) tends to increase the time animals spent awake in lateral recumbency in direct sunlight, or in the authors’ terms, ‘sun-basking’ (Hogan et al., 2010). Moreover, evolutionary perspectives predict that behaviours that help maintain homeostasis and promote evolutionary fitness are likely to often produce rewarding sensations (Cabanac, 1971; Fraser and Duncan, 1998). Tawny frogmouths (*Padargus strigoides*; Kortner and Geiser, 1999) and diurnal striped mice (*Rhabdomys pumilio*; Schradin et al., 2007) do chose to stand motionless or lie awake in

direct sunlight in cold conditions, a so-called ‘sun-basking’ behaviour that presumably helps maintain homeostasis by facilitating passive thermoregulation and removes the aversive feeling of coldness. Similarly, poikilothermic animals actively chose to sun-bask until their body temperatures reach their preferred body levels (*e.g.* in turtle *Pseudemys Scripta*, Crawford et al., 1983; in Nile crocodile *Crocodylus niloticus*, Downs et al., 2008; in blue spiny lizard *Sceloporus cyanogeny*, Garrick, 1979). Interestingly, Fell et al. (2014) have recently shown that chronic low doses of UV exposure elevate laboratory mice plasma levels of β -endorphin, an endogenous opioid known to play a role in reinforcement. While one may question the biological relevance of UV exposure in a nocturnal animal, these results nevertheless suggest that sun-basking could have biologically relevant rewarding properties in diurnal rodent species, such as the above-mentioned striped mouse (Schradin et al., 2007). Thus, although the evidence is not yet conclusive, additional empirical tests of the hypothesis that sun-basking is pleasurable in animals, as in humans, seem very worthwhile (*e.g.* in domestic cats, anecdotally reported by their owners to lie down in certain areas at the time of the day these areas are sunny: Fureix, personal observation).

4.2.2. Post-consummatory inactivity

Post-consummatory inactivity, such as inactivity immediately expressed post-copulation, is likely to be associated with satisfaction and to be pleasurable. In humans, experiencing sexual arousal to orgasm usually produces a pleasant calming effect of sexual satisfaction (Graber et al., 1985; Levin, 2007), and partners frequently remain relatively inactive during the post-coital time (*e.g.* remaining awake and cuddling with the partner, or falling asleep, Hughes and Kruger, 2011). Remaining inactive close to their mates after copulation has also been reported in animals. For instance, in horses, immediately following ejaculation, the stallion’s body relaxes, and its head droops beside the mare’s neck for a few seconds, after

which the stallion dismounts and commonly stands quietly behind the mare, often relaxed and inactive (Waring, 2003, P168). Rams similarly usually remain standing quietly beside the female with their heads down slightly shortly after ejaculation and dismounting (Pepelko and Clegg, 1965), while some mink remain motionless, as if sunk in deep stupor, for a period of time after mating (Diez-Leon, 2014, personal communication). Mating mice generally fall over onto their sides for 5-10 seconds immediately post-ejaculation whilst still coupled, with open eyes and apparent unresponsiveness to sensory stimuli (*e.g.* being touched with a finger) (Brennan, 2015, personal communication). Following evolutionary predictions (*e.g.* Cabanac, 1971; Fraser and Duncan, 1998) and by analogy with humans (Graber et al., 1985; Levin, 2007), sexual interaction and orgasm are typically likely to be pleasurable (see also Dixon, 2010, P392-393), at least in healthy male mammals (in which ejaculation can be observed). Due to its very close temporal relationship with the sexual interaction, one may reasonably hypothesize that this post-copulation inactivity in animals (or, at least, in male mammals) has, just as in humans, a pleasant ‘calming’ affective component (see 4.2.3 and discussion for further research suggestions).

Inactivity expressed in postprandial contexts could also be associated with positive affective states. Postprandial inactivity is likely to be associated with satiety, *i.e.* in humans the feeling of ‘fullness’ following a feeding episode (Benelam, 2009; Harrold et al., 2012), and has been observed in a number of animal species (rats: *e.g.* Richter, 1922; Antin et al., 1975; Willner et al., 1990; Rodgers et al., 2010; northern harriers *Circus cyaneus*: Temeles, 1989; sows: Zonderland et al., 2004; dogs: Bosch et al., 2009; cats: Fara and colleagues 1969, cited in Orr et al., 1997). Further evidence that postprandial inactivity in animals is likely to be associated with satiety comes from pharmacological studies: cholecystokinin⁵ administration, which in humans increases the feeling of fullness and reduces food

⁵ a group of peptides localized in the gut in mammals

consumption (*e.g.* Stacher et al., 1979; Crawley et al., 1982; Stacher et al., 1982; Sam et al., 2012), also reduces food consumption in animals (rats, mice, sheep, pigs, monkeys, reviewed in Crawley et al., 1982), and induces inactivity (rats: *e.g.* Antin et al., 1975; mice: Crawley et al., 1981; rhesus monkeys: Falasco et al., 1979). In calves, being able to suck on a teat increases the tendency to rest after milk consumption (Veissier et al., 2002), and this may be mediated in part by cholecystokinin, which increases in response to such sucking (De Passillé et al., 1993).

Satiety *can* increase inactivity in humans as well (*e.g.* napping after lunch, Zammit et al., 1992; Vela-Bueno et al., 2008), although the causal relationship remains debated (review in Campbell, 1992), making it difficult to use evidence regarding affective states during satiety-induced inactivity in humans. Nevertheless, because satiated humans self-report positive affective states, such as satisfaction and relaxation (Panksepp, 2005; Boelsma et al., 2010; Seehuus et al., 2013), one may reasonably hypothesize that postprandial inactivity in animals has a positive affective component just as humans experience after eating. This suggestion is supported by the finding that, in laying hen chicks, denying access to the part of a pen designed to accommodate postprandial inactivity results in a more negative affective state than in a baseline situation (where chicks have free access to that area), as evidenced by a more ‘pessimistic-like’ response in a judgment bias paradigm (Seehuus et al., 2013).

With respect to *satiation*, *i.e.* the *processes* that bring episodes of eating behaviour to an end (Benelam, 2009; Harrold et al., 2012), ingesting food is typically considered activity in applied ethology research (see section 2); satiation therefore appears unlikely to involve inactivity in most species, including humans. However, rumination, which has the primary function of facilitating clearance of digesta from the rumen by reduction of particle size, is most frequently expressed when the animals are motionless lying down (Wagnon, 1963; Kilgour, 2012; Schirmann et al., 2012). As such, it could be seen as a (ruminant-specific)

657 satiation-induced specific form of inactivity. In humans, satiation is associated with positive
658 affective states, such as feelings of liking and satisfaction (*e.g.* Benelam, 2009; Seehuus et al.,
659 2013), and lying down ruminating has been suggested to be ‘a sign of relaxation in cattle’
660 (Phillips, 2002, from Espejo and Endres, 2007) and a sign that cows are ‘at ease’ (Bristow
661 and Holmes, 2007). Rumination also appears to decrease when animals are exposed to
662 aversive situations, such as social stressors (regrouping) and home-pen novelty (Schirmann et
663 al., 2011), disturbance by flies (presumably associated with discomfort: Wagnon, 1963, p47)
664 and ruminal acidosis (presumably associated with pain, lactating dairy cows: DeVries et al.,
665 2009).

667 4.2.3. Inactivity and positive affective states: further research suggestions

668 Being *inactive* in order to *favour* reward *acquisition* is likely rare, with perhaps the only
669 direct evidence coming from humans during yogi meditation, as discussed above. Our other
670 suggested example, bouts of immobility while stalking in cats, is clearly not validated as an
671 indicator of positive affective states in animals. Potentially more fruitful as affective state
672 indicators are specific forms of inactivity reported in humans and expressed in animals when
673 threat levels are low, such as sun-basking and post-consummatory inactivity. All of these still
674 need validation, however. Further work could investigate to what extent acute stressors or
675 chronically aversive environments would decrease such inactivity (and the opposite for
676 preferred environments), how these forms of inactivity would be modulated by
677 pharmacological manipulations inducing either negative or positive affective states, and,
678 more specifically, whether the putatively positive affective state associated with postprandial
679 inactivity would be lessened in force-fed animals (see Faure et al., 2001 for evidence that
680 force-feeding might be perceived as aversive in ducks). Further research is also warranted
681 into other forms of inactivity likely to be pleasurable in humans, such as being passively

rocked and breast-feeding, which could perhaps find equivalents in animals *choosing* to go and float in the water and in lactating females.

5. Misinterpreting inactivity as an indicator of affective state

We will further discuss the relationship between increased inactivity and affective states by presenting misleading situations prone to yield wrong conclusions. First, we will describe some examples of ‘false negatives’ (*i.e.* those cases where the animal’s affective state is likely to be either poor or good but the animal *is not* inactive), and ‘false positives’ (*i.e.* those cases where the animal *is* inactive, eliciting interpretations about its affective state, while the animal actually *does not* experience the presumed affective state). We will then discuss specific forms of inactivity which appear to be not necessarily linked to an actual affective state, but instead to a *lack* of emotion, and how the methodologies used to assess inactivity could yield different interpretations with regards to its associated affective states, discussing in detail the case of resting.

5.1. The risk of wrong conclusions: some examples of ‘false negatives’

As discussed above (sections 3 and 4), specific forms of inactivity and/or overall decreases in activity in many contexts are likely to be associated with specific affective states; as such, an inactive animal in a similar situation is believed to be in a more intense (negative or positive) affective state than its comparatively more active counterparts. Assuming this *systematically* would nevertheless sometimes yield incorrect conclusions; in some situations, animals are likely to *experience the* specific (negative or positive) *affective states* of interest, but *do not display* increased inactivity.

Inactivity can sometimes be one of two (or more) alternative responses – driven by individual characteristics – to the same situation, which both indicate a similarly (in this

example) negative affective state. For instance, while individuals can respond to situations involving a (perceived) threat by freezing or hiding, they can also display active reactions, such as fleeing or even attacking (Boissy, 1995; Blanchard et al., 2011). In red deer, the response strategies to a perceived threat differ according to age: juveniles employ a hiding strategy, and freeze in response to threat, but as they age, they begin fleeing from some threats instead (Espmark and Langvatn, 1985). This does not mean we should conclude that adult red deer are not afraid of the perceived threat from which they are fleeing only because they are not displaying inactive responses. More generally, personality can determine the form of an individual's response, including whether they become inactive or not. For example, speed of exploration of a novel environment is considered to reflect a personality trait in birds, and individuals with different exploratory phenotypes also differ in the degree to which they become inactive after social defeat (reviewed by Groothuis and Carere 2007). Van Reenen et al. (2005) thus suggest that their failure to find a correlation between open field locomotion and other measures of response to novelty might be explained by the presence of different coping styles, such that some calves responded to the open field with escape attempts, but others with immobility, novelty being nevertheless perceived as frightening in both cases. Moreover, an individual's experience is likely to influence its stress responses: for example, captive-born individuals may be more likely to respond actively to a sub-optimal captive environment, developing stereotypic behaviour, while wild-caught individuals may be more likely to respond by hiding (*e.g.* Jones et al., 2011; Camus et al., 2013). Despite these differences, individuals displaying both response types are likely to suffer from their sub-optimal life conditions.

Inactivity can also be one of two (or more) alternative responses – dependent this time on situational characteristics - to different situations associated with a similar (in this example negative) affective state. For instance, Cooper et al. (1996) showed that voles responded to an

732 unfamiliar sound by freezing if in an enriched environment where cover was available, but
733 otherwise responded actively, by running or digging; in such a case, there is no good reason
734 to conclude that running or digging animals are not afraid of the unfamiliar sound simply
735 because they do not freeze. Moreover, while ill-health (including painful) conditions increase
736 inactivity in a number of species, including humans (see 3.2.a.), both increased and decreased
737 sleep are used by caregivers as behavioural signs of pain in non-verbal cognitively impaired
738 children (McGrath et al., 1998), and animals sometimes also display active behaviours in
739 response to ill-health conditions. For instance, in mice experiencing scrotal approach
740 vasectomy, Leach et al. (2012) observed higher frequencies of pain behaviours (*e.g.* circle,
741 flinch, stagger, twitch and writhe) and higher Mouse Grimace Scale (MGS) scores in the
742 animals receiving a saline solution post-operatively, compared to pre-surgery periods and to
743 mice receiving post-operative analgesia (meloxicam, bupivacaine). Mice without post-
744 operative analgesia are likely to experience pain; however, none of the inactive behaviours
745 recorded in the study (*e.g.* ‘stand’ and ‘sleep’) differed pre- vs. post-surgery, nor between
746 treatment groups. One may hypothesize that at least some of the active pain-related
747 behaviours might allow animals to cope better with pain induced by the surgery than being
748 inactive. Although this hypothesis remains to be tested in the context of that study, focusing
749 only on the absence of increased inactivity here would lead to the conclusion that the mice
750 experiencing scrotal approach vasectomy without post-operative analgesia do not suffer,
751 which is contradicted by the displayed pain behaviours and MGS scores. Another example
752 comes from a study on Pekin ducks (*Anas platyrhynchos*), in which animals were provided
753 with environmental options allowing them to actively attempt to cope with the situation.
754 When injected either with saline solution or pathogen-associated molecular patterns, saline-
755 injected ducks exhibited pronounced anorexia strongly correlated with a fever response, but
756 none of the treatments significantly affected the level of animals’ activity, measured by

activity loggers surgically inserted into the abdominal cavity (Marais et al., 2013). According to the authors, sick ducks might have actively attempted to lower their body temperature during the defervescent phase of fever by getting in and out of the bathing tub provided. This behaviour, which the authors had previously observed in ducks given pyrogens, would have contributed to the amount of activity logged on the days when ducks were given pathogen-associated molecular patterns.

5.2. *The risk of wrong conclusions: some examples of ‘false positives’*

Erroneous interpretations might also come from those cases of ‘false positives’, where an animal *is* inactive, raising interpretations about its affective state, while the animal’s actual affective state *does not differ*, or *even goes the opposite direction*, from the affective state of its comparatively more active counterparts. For instance, if animals are afraid or motivated to hide but unable to do so because no appropriate camouflaged hiding places are available, their welfare is not likely to be better than if they were hiding (*e.g.* leopard cats: Carlstead et al., 1993a; Wielebnowski et al., 2002; mink: Nimon and Broom, 1999; shelter cats: Kry and Casey, 2007). Or, in other words, if animals are afraid or motivated to hide and able to do so because the cage provides them with a hiding place, there is no good reason to conclude that because they are inactive, their welfare is worse than the welfare of their counterparts who are in the same situation but prevented from hiding. Similarly, successful environmental enrichment often decreases inactivity in a wide range of species (*e.g.* Anna et al., 2002; Koistinen et al., 2009; Rozek et al., 2010), with the exceptions to this rule being types of physical enrichment that would primarily be expected to increase comfort or perceived safety, such as shelters (Wurbel et al., 1998a; Tilly et al., 2010); it would be absurd to conclude that providing animals with shelters decreases their welfare.

While the examples above highlight cases of false positives when inactivity is expected to be associated with negative affective states, false positives can also happen when inactivity is expected to be associated with positive affective states. For instance, Mason and Latham (2004) found in a meta-analysis that stereotypic behaviours are more prevalent in populations living in sub-optimal conditions than in populations kept under more welfare-friendly conditions, but also that, more often than not, within populations where stereotypic behaviour was prevalent, individuals that did not stereotype, or had relatively low levels of stereotypy, had poorer welfare than those that with high levels, according to a variety of welfare measures. Since non-stereotypic individuals are likely to be the most inactive individuals within a population (*e.g.* Bildsoe et al., 1990; Wurbel et al., 1998b), this may indicate that at an individual level, inactive responses to stressful conditions are actually more often associated with poor welfare than with good welfare.

5.3. *The risk of wrong conclusions: cases where inactivity is linked to a lack of emotion*

While the examples above discuss specific forms of inactivity which are likely to be associated with (either negative or positive) affective states, some forms of inactivity appear to be not necessarily linked to an actual affective state but instead a *lack* of emotion. For instance, disorders of reduced motivation such as apathy, defined as ‘a state of diminished motivation in the presence of normal consciousness, attention, cognitive capacity, and mood’ (Marin and Wilkosz, 2005)⁶ involve decreased activity. In Marin and Wilkocz’s words (our emphasis), ‘patients with diminished motivation **all show diminished activity**’; however they are also ‘**emotionally indifferent**... or display restricted responses to important life events’. Another example is a ‘deconstructed state’ observed in the pre-suicidal phase and in

⁶ Apathy is not the only disorder of diminished motivation, but identifying the two other common disorders (abulia and akinetic mutism) relies on speech; therefore those states are not currently possible to operationalise in non-humans and of less interest here.

socially excluded individuals, defined as a ‘defensive state of cognitive deconstruction that avoids meaningful thought, **emotions** and self-awareness, and **is characterized by lethargy and passivity** and alerted time flow’ (cited from Twenge et al., 2003, emphasis ours). In animals, Engel and Schmale (1972) described a broad category of stress-induced forms of inactivity that include decreased responsiveness to the environment, can persist over a long period of time, and are believed to be adaptive because they reduce predation risk and allow the conservation of significant amounts of energy; they called this category conservation-withdrawal (C-W). The actual valence of the affective state associated with C-W, if any, is still debated, with some authors describing it as an ‘affectively neutral’ state (Weiner and Lovitt, 1979).

While these forms of inactivity have (or, in the case of C-W, could have) no affective component at the time they are displayed, it seems worth noting that they all appear in negatively valenced contexts, and in humans, often yield situations from which individuals are likely to suffer, such as conflicts with relatives due to family burden. Therefore, even though these specific forms of inactivity cannot be considered as *indicators* of the individual’s actual affective state, they should nevertheless be taken as a sign of exposure to suboptimal environments, and potential poor welfare.

5.4. *Different methodologies, different conclusions?*

As previously highlighted (section 2), there is some variation in the literature in exactly what the term ‘inactivity’ encompasses. While how inactivity is assessed depends on one’s perspective and hypotheses under test, methodological variation in terms of how behaviour is categorized as inactive *versus* active is likely to yield some diversity in the effects observed in terms of welfare states associated to inactivity. For instance, relying only on pedometers, radio-tracking or similar technology -- where any time not in locomotion is called inactivity --

would not discriminate *e.g.* a motionless healthy animal sun-basking (likely to experience positive affective states, see 4.2.1) from an animal awake but inactive due to injury (likely to experience negative affective states, see 3.2.1.).

A detailed example of how methodological variations in defining inactivity could influence its interpretation in terms of associated affective states comes from resting. Resting can be seen as a post-consummatory (of various activities) behaviour, and is often considered to reflect positive affective states. Indeed, safe, comfortable contexts promote rest (*e.g.* larger home stalls in horses: Raabymagle and Ladewig, 2006). Preferred situations often decrease signs of poor welfare and increase time resting (enriched cages, in rats: Abou-Ismaïl and Mahboub, 2011; and in mice: Tilly et al., 2010; bedding types that are preferred when the animals are given a choice of stalls in horses: Hunter and Houpt, 1989; Mills et al., 2000; Pedersen et al., 2004; Werhahn et al., 2010). So does providing a more naturalistic social environment in horses by introducing adult conspecifics in groups of sub-adults (Bourjade et al., 2008). Moreover, a variety of stressors, such as chronic exposure to mild unpredictable stressors (rats: Cheeta et al., 1997), exposure to an aggressive dominant conspecific (male tree shrews *Tupaia belangeri*: Fuchs and Flugge, 2002) and social isolation (rats: Hurst et al., 1999), decrease the time the animals spend resting over hours or days. Rats exposed to sleep disturbance (husbandry procedures performed during the non-active light phase) not surprisingly sleep less, spending more time *awake non-active* and show higher indicators of physiological stress and reduced welfare than do their conspecifics experiencing husbandry procedures during their active dark phase (Abou-Ismaïl et al., 2008). Moreover, while provoking sexual (positive) interactions during the inactive phase only briefly suppresses sleep in male mice, aversive social conflict induces 12h long-lasting sleep disturbances (Meerlo and Turek, 2001). Resting is therefore commonly interpreted as a sign that animals are relaxed and experience positive affective states.

However, in a number of animal studies, resting and *sleeping* are merged together in the behavioural repertoire, defined by the animal displaying a species-specific posture (usually lying down, but sometimes also sitting or even standing still) *with eyes partially or fully closed* (e.g. in rats: Hurst et al., 1999; Abou-Ismaïl et al., 2007; 2008; in rabbits: Zeidner et al., 1983; in birds: Campbell and Tobler, 1984; in horses: Waring, 2003). This is because measuring sleep, a restorative behaviour ‘not distinguished by movement’ (Carlson, 2012, p289) and which ‘can be defined behaviourally by the normal suspension of consciousness and electrophysiologically by specific brain wave criteria’ (Purves et al., 2007, p 707) requires performing invasive and/or technically challenging (in animals) measurements, *i.e.* electroencephalographic (EEG) measurement together with electromyographic (EMG) signals. Following the rationale that the longer a bout of inactivity, the more likely it is to be sleep, some authors have investigated duration of inactivity as a way to estimate sleep in mice (Pack et al., 2007). While inactivity-defined sleep (in that study being motionless for ≥ 40 s) and EEG-EMG defined sleep did show good convergence in mice (Pack et al., 2007), using such a duration-of-inactivity-only criterion to estimate sleep appears however to be prone to yield false positives. It would not discriminate, for instance, a healthy animal sleeping from an animal awake but inactive due to pain (see part 3). While EEG-EMG therefore appear to remain the ‘gold standard’ methods to measure sleep, using such techniques is however often too challenging (practically) to perform in most of the farm, zoo and companion animal species, and therefore cannot be applied in a vast number of studies in applied ethology research.

Despite (realistically good) reasons to do so, merging sleep and rest in the animals’ behavioural repertoire could have significant implications with respect to the welfare states associated to this inactivity. Indeed, in humans, neither sleep quantity nor its quality appear to

879 be trustworthy indicators of affective states. For instance, insomnia⁷ (APA, 2013) can be
880 caused either by negative feelings such as stress or pain (Purves et al., 2007, p728; Carlson,
881 2012, p298), or by ‘excited anticipation of a pleasurable event’ (Carlson, 2012, p298). Sleep
882 disturbance is also a prominent symptom of clinical depression (estimated to affect up to 90%
883 of those with depression, Paterson et al., 2009), but either insomnia or hypersomnia can be
884 observed (*e.g.* APA, 2013; WHO, 1994; Maurice-Tison et al., 1998; Henn and Vollmayr,
885 2005), with some depressed people even self-reporting mixed insomnia/hypersomnia
886 symptoms (Paterson et al., 2009). Similarly, both insomnia and hypersomnia are part of the
887 diagnostic symptoms under ICD-10 for withdrawal states from stimulants, with people
888 experiencing such states also self-reporting negative emotions such as ‘depressed mood’, and
889 ‘decreased contentedness / well-being’ (Juliano and Griffiths, 2004). It is also worth noting
890 that sleep manipulations yield quite erratic effects on people’s mood. In healthy people, acute
891 and short-term sleep deprivation usually worsens mood (*e.g.* Weinger and Ancoli-Israel,
892 2002; Drury et al., 2012), but extending sleep (*e.g.* by 2 or 3h per night beyond its habitual
893 duration) has been reported to worsen mood, to improve mood or to induce no mood change
894 (David et al., 1991; Ferrara and De Gennaro, 2001). Moreover, while sleep deprivation in
895 healthy people usually worsens mood, it is usually followed by a short-term mood
896 *improvement* in depressed patients (Benedetti and Colombo, 2011).

897 Thus, while a number of studies in animals do support the view that ‘resting’ is likely to
898 be associated with positive affective state (enhanced in preferred / positive contexts, reduced
899 in aversive conditions), it seems worth noting that, in the majority of these studies, the
900 relative proportion of the observed inactivity that is ‘simply’ resting cannot be disentangled
901 from that ‘purely’ sleeping due to methodological challenges. As human studies show that
902 sleep is clearly not a trustworthy indicator of the affective state (its duration and quality can

⁷ ‘subjective complaint of difficulty falling asleep or staying asleep, or poor sleep quality’, DSM-V p823

be modified in either direction under either positive or negative affective states), the interpretation of the ‘resting + sleeping’ behaviour in animals in terms of its associated affective state might not be so straightforward.

6. Further research directions

Can inactivity -- in its various forms – be a useful indicator of specific affective states in animals? We think it can, based on analogies with humans self-reporting their feelings while displaying specific forms of inactivity phenotypically similar to those displayed by animals in similar situations (summarised in Table 1). Most of the specific forms we discussed in this paper still need further refinement and validation before they could be used in this way, however.

While some forms are unambiguously specific behaviours (*e.g.* freezing in rodents, ‘withdrawn’ states in horses) or operationally definable (*e.g.* with regards to location or other activities such as immediately post-mating) (see column 4 in Table 1), others are less specifically described, such as overall increased inactivity / decreased activity in inactive ill animals or individuals displaying signs of learned helplessness. Reassuringly, contextual information probably favours correct recognition of the associated affective state: an unmedicated animal being inactive post-surgery or displaying signs of ill-health (*e.g.* fever, injury) is likely to experience the aversive affective component of ill-health conditions, while exposure to inescapable, uncontrollable stressors is unlikely to induce positive feelings. Given such contextual knowledge, inactivity may be useful as an indicator of intensity of the affective state. Effort should nevertheless be made in the future to define these specific forms of inactivity more precisely if relevant to hypothesis under test (*e.g.* by adding fine postural descriptions) (cf. also 5.4.).

Further work could also investigate how those specific forms of inactivity that are currently not pharmacologically validated (see Table 1, last column) would be modulated by giving the animals drugs inducing either negative or positive affective states. This is provided that such drugs have already been validated as inducing the affective state of interest for the tested species, and are known *not* to induce sedative side effects (risk of circular reasoning otherwise). Such validation would be of primary interest for any forms, but particularly for those few forms observed in animals which are obviously not (food rumination) or not systematically (satiety-induced inactivity) displayed by people, making it difficult in these few cases to use humans-based evidence to infer the associated affective states in animals.

Beyond this, future validation work could investigate to what extent acute stressors and chronically aversive environments increase those specific forms of inactivity believed to reflect negative affective states (with the opposite being the case for preferred environments), to provide a starting point for discriminating between forms of inactivity reflecting short- and/or long-term affective states. It could also investigate the co-variation of a specific form of inactivity with evolutionary fitness (following evolutionary perspectives that predict that individuals are likely to avoid aversive sensations and pursue rewarding sensations that respectively decrease and promote evolutionary fitness, Cabanac, 1971; Fraser and Duncan, 1998); and could investigate whether a specific form of inactivity co-varies with other welfare indicators, provided these are previously-validated indicators of the specific affective state under test (for instance excluding cortisol levels, reported to either increase or decrease in chronically stressed individuals as well as to increase in some positive situations, *e.g.* in humans: Miller et al., 2007, in animals: Rushen, 1991; Mormede et al., 2007).

7. Conclusions

952 Should more attention be paid to inactivity in behavioural and animal welfare studies?
953 Considering subtypes of inactivity, we think it should. First, as discussed in this paper, some
954 specific forms of inactivity (*e.g.* displayed in response to a perceived threat) are useful
955 indicators of poor welfare states. A number of others forms have, acknowledging that further
956 refinement and validation are still needed, the potential to indicate either negative or positive
957 affective states in animals. This makes additional research into this topic very worthwhile.
958 Moreover, even when inactivity does not result from poor welfare, levels of inactivity that are
959 too high or too low can directly or indirectly induce poorer affective states, raising welfare
960 concerns. For example, in group-housed hens, inactive hens are more likely to be victims of
961 feather pecking, and thus suffer due to high inactivity levels (Riber and Forkman, 2007).
962 Meanwhile, retaining adaptive forms of inactivity is still essential: in reintroduction
963 programmes for endangered species, individuals that have not been active enough to learn
964 appropriate skills before being reintroduced into the wild are likely to have poor welfare once
965 released, since they may be unable to attain sufficient food or find shelter, and may be at
966 higher risk of injury (McPhee, 2004).

967 Using inactivity as an indicator of affective states in animals does require a number of
968 changes in the way we often view inactivity in behavioural and animal welfare studies,
969 however. As discussed throughout this paper, inactivity is *not* a homogeneous category of
970 behaviour: there are many different, context- (and sometimes species-) specific forms.
971 Merging these specific forms into a single broad category certainly can yield erroneous
972 interpretations with regards to the associated affective states, by *e.g.* not discriminating a
973 healthy animals resting from an animal inactive due to ill-health conditions. Prior to data
974 collection, ethograms should include precise descriptions (*e.g.* by adding fine postural
975 descriptions) of any specific form(s) of inactivity relevant to hypotheses under test. A clear
976 description of which contexts trigger (or conversely, decrease) specific form(s) of inactivity

is also crucial, as it is the first, essential, step towards inferring its putatively associated affective state(s). Additional justifications should be provided before inferring putative affective states associated with inactivity in animals. Bringing human and animal studies together to rely on analogies with humans self-reporting their feelings is one of the possible justifications; so are pharmacological approaches, which strengthen construct validity. Further research suggestions mentioned in this paper would certainly deepen our understanding of what inactivity can reveal about affective states in non-human animals, providing new ways of assessing treatment effects and a better understanding of the implications of personality differences.

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1666

1667 **10. Table legend**

1668 Table 1. Specific forms of inactivity and their association with specific affective states.

Table

Name	Valence	Arousal	Specific form of inactivity, defined operationally	Expressed in response to:	In humans this situation (or similar) has been reported to be:	Humans display a phenotypically similar inactivity:	Pharmacological evidence?
Freezing	Negative	High	Yes	(Perceived) actual or potential threat	Aversive	Yes	Yes (enhanced by anxiogenics, reduced by anxiolytics)
Tonic immobility	Negative	High	Yes	(Perceived) actual or potential threat	Aversive	Yes	Yes (enhanced by anxiogenics, reduced by anxiolytics)
Hiding	Negative	High	Yes (provided hiding opportunities)	(Perceived) actual or potential threat	Aversive	Yes	Yes (enhanced by anxiogenics, reduced by anxiolytics)
Ill-health inactivity	Negative	Low	No (decreased activity)	Illness, injury	Aversive	Yes	Yes (reduced by analgesics)
Learned helplessness related ‘floating in despair’ (laboratory rodents)	Negative	?	Yes	Porsolt (forced swim) test	Aversive	N/A (specific testing conditions in rodents are not transferable to humans)	Yes (reduced by antidepressants)
Learned helplessness related overall passivity	Negative	Low	No (decreased activity)	Inescapable / uncontrollable aversive environments	Aversive	Yes	?
Depression-like ‘withdrawn’ state (horses)	Negative	Low	Yes	?	N/A	No phenotypically exactly similar form	?
Boredom-like lying down with eyes open (mink)	Negative	Low (but debated)	Yes	Barren, impoverished environments	Aversive	Yes, but restlessness may also be seen	?
Standing/lying in the sun	Positive	Low	Yes (provided chose shaded / sunny areas)	Sunny area	Positive	Yes	Yes (induces a β -endorphin release)
Post-copulation inactivity	Positive	Low	Yes (timing mating)	Immediately after mating	Positive	Yes	?
Satiety-related inactivity	Positive	Low	Yes (timing eating)	Post-prandial	Positive	Unsure	?
Rumination (cattle)	Positive	Low	Yes	Post food consumption	N/A	N/A	?
Stillness when stalking (cats)	Positive	High	Yes	Predation, play	?	?	?